

Food and Drug Administration 10903 New Hampshire Avenue Document Control Center – WO66-G609 Silver Spring, MD 20993-0002

December 31, 2014

Gyrus ACMI, Incorporated Mr. Neil Kelly MBA, RAC Senior Regulatory Affairs Specialist 6655 Wedgwood Road Maple Grove, Minnesota 55311

Re: K142289

Trade/Device Name: PK Spatula Regulation Number: 21 CFR 878.4400

Regulation Name: Electrosurgical cutting and coagulation device and accessories

Regulatory Class: Class II

Product Code: GEI

Dated: December 4, 2014 Received: December 5, 2014

#### Dear Mr. Kelly:

We have reviewed your Section 510(k) premarket notification of intent to market the device referenced above and have determined the device is substantially equivalent (for the indications for use stated in the enclosure) to legally marketed predicate devices marketed in interstate commerce prior to May 28, 1976, the enactment date of the Medical Device Amendments, or to devices that have been reclassified in accordance with the provisions of the Federal Food, Drug, and Cosmetic Act (Act) that do not require approval of a premarket approval application (PMA). You may, therefore, market the device, subject to the general controls provisions of the Act. The general controls provisions of the Act include requirements for annual registration, listing of devices, good manufacturing practice, labeling, and prohibitions against misbranding and adulteration. Please note: CDRH does not evaluate information related to contract liability warranties. We remind you; however, that device labeling must be truthful and not misleading.

If your device is classified (see above) into either class II (Special Controls) or class III (PMA), it may be subject to additional controls. Existing major regulations affecting your device can be found in the Code of Federal Regulations, Title 21, Parts 800 to 898. In addition, FDA may publish further announcements concerning your device in the <u>Federal Register</u>.

Please be advised that FDA's issuance of a substantial equivalence determination does not mean that FDA has made a determination that your device complies with other requirements of the Act or any Federal statutes and regulations administered by other Federal agencies. You must comply with all the Act's requirements, including, but not limited to: registration and listing (21 CFR Part 807); labeling (21 CFR Part 801); medical device reporting (reporting of medical

device-related adverse events) (21 CFR 803); good manufacturing practice requirements as set forth in the quality systems (QS) regulation (21 CFR Part 820); and if applicable, the electronic product radiation control provisions (Sections 531-542 of the Act); 21 CFR 1000-1050.

If you desire specific advice for your device on our labeling regulation (21 CFR Part 801), please contact the Division of Industry and Consumer Education at its toll-free number (800) 638-2041 or (301) 796-7100 or at its Internet address

http://www.fda.gov/MedicalDevices/ResourcesforYou/Industry/default.htm. Also, please note the regulation entitled, "Misbranding by reference to premarket notification" (21CFR Part 807.97). For questions regarding the reporting of adverse events under the MDR regulation (21 CFR Part 803), please go to

http://www.fda.gov/MedicalDevices/Safety/ReportaProblem/default.htm for the CDRH's Office of Surveillance and Biometrics/Division of Postmarket Surveillance.

You may obtain other general information on your responsibilities under the Act from the Division of Industry and Consumer Education at its toll-free number (800) 638-2041 or (301) 796-7100 or at its Internet address

http://www.fda.gov/MedicalDevices/ResourcesforYou/Industry/default.htm.

Sincerely yours,

# David Krause -S

for Binita S. Ashar, M.D., M.B.A., F.A.C.S.
Director
Division of Surgical Devices
Office of Device Evaluation
Center for Devices and
Radiological Health

Enclosure

# DEPARTMENT OF HEALTH AND HUMAN SERVICES

Form Approved: OMB No. 0910-0120

Food and Drug Administration	Expiration Date: January 31, 2017
Indications for Use	See PRA Statement below.
10(k) Number (if known)	
TBD K142289	
Device Name	
PK Spatula	
ndications for Use <i>(Describe)</i> The PK Spatula is indicated for resection and coagulation of sof	it issue and blood vessels in lanaroscopic and general
surgical procedures when used with the Olympus ESG-400 Gen	
The of the (Colori and as both as applicable)	
Type of Use (Select one or both, as applicable)	
☑ Prescription Use (Part 21 CFR 801 Subpart D)	Over-The-Counter Use (21 CFR 801 Subpart C)
PLEASE DO NOT WRITE BELOW THIS LINE - CO	NTINUE ON A SEPARATE PAGE IF NEEDED.
FOR FDA US	SE ONLY
Concurrence of Center for Devices and Radiological Health (CDRH) (S	Signature)
Concurrence of Center for Devices and Radiological Health (CDRH) (S	Signature)

This section applies only to requirements of the Paperwork Reduction Act of 1995.

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> Department of Health and Human Services Food and Drug Administration Office of Chief Information Officer Paperwork Reduction Act (PRA) Staff PRAStaff@fda.hhs.gov

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FORM FDA 3881 (1/14) Page 1 of 1 PSC Publishing Services (301) 443-6740 EF Olympus PK Spatula Gyrus ACMI, Inc. Traditional 510(k) Notification August 15, 2014

K142289

# 510(k) Summary of Safety and Effectiveness Gyrus ACMI, Inc.

# Olympus PK Spatula

# **General Information**

Manufacturer: Gyrus ACMI, Inc.

6655 Wedgwood Road Maple Grove, MN 55311 Phone: 508-804-2600

Establishment Registration Number: 2183680

510(k) Submitter: Gyrus ACMI, Inc.

136 Turnpike Rd.

Southborough, MA 01772-2104

Establishment Registration Number: 3003790304

Contact Person: Neil Kelly

Regulatory Affairs Specialist

508-804-2690

Neil.kelly@olympus-osta.com

Date Prepared: August 15, 2014

**Device Description** 

Classification Name: Electrosurgical cutting and coagulation

device and accessories

Regulation Number 21 CFR 878.4400

Product Code GEI Regulatory Class II

Review Panel General and Plastic Surgery

Trade Name: PK Spatula

Generic/Common Name: Spatula Electrode

# **Predicate Devices**

Gyrus PlasmaCision Laparoscopic Spatula K041633

Traditional 510(k) Notification August 15, 2014

#### **Comparison to Predicate Devices:**

The PK Spatula has been compared to our own legally marketed Gyrus PlasmaCision Laparoscopic Spatula (K041633) with respect to intended use and technological characteristics. The comparison and testing results presented in this 510(k) notification to the FDA show that the device is substantially equivalent to predicate devices and raises no new concerns or safety or effectiveness.

#### **Product Description**

The PK Spatula is a bipolar electrosurgical instrument with the capability to cut and coagulate soft tissue and blood vessels in laparoscopic and general surgery. The instrument will pass through a 5mm cannula or through an operating laparoscope working channel of 5mm or larger diameter. The instrument is to be used only with the ESG-400 electrosurgical generator. The generator and device make up a medical electrical system. The instrument is to be used only with the Gyrus ESG-400 Generator and associated 5 way connector cable. The device is intended for use in a non-irrigated (dry) environment.

## **Technological Characteristics**

The PK Spatula uses bipolar energy in order to cut and coagulate soft tissue and blood vessels in laparoscopic and general surgical procedures. The PK Spatula is activated using buttons located on the device handle. This allows the physician to activate either cut or coagulation (coag) mode without taking their eyes off the surgical site. Historically foot pedals have been used for such devices and are also available for the proposed device. A nosecone located at the distal end of the handle and at the proximal end of the device shaft allows the physician to alter the orientation of the electrode tip without altering the orientation of the handle.

#### Material

The predicate and proposed devices share many common materials. The two patient contact material differences are the sheath, which is now flouropolymer rather than Polyimide tubing, and a new ink was added on the device shaft as well. Biocompatibility testing has been carried out with passing results. As for the electrode tip and insulation all materials remain the same as the predicate.

#### **Intended Uses**

The PK Spatula is indicated for resection and coagulation of soft tissue and blood vessels in laparoscopic and general surgical procedures when used with the Olympus ESG-400 Generator.

Olympus PK Spatula Gyrus ACMI, Inc.

#### **Compliance to Voluntary Standards**

The design of the proposed device complies with the following standards:

ISO 10993-1, 2009 ISO 10993-5, 2009 ISO 10993-7 2008 ISO 10993-10, 2010 ANSI/AAMI/ISO 11607-1, 2006 ANSI/AAMI/ISO 11135-1, 2007 ISO 14971, 2007 ISO 15223-1; 2012 IEC 60601-1: 2005 IEC 60601-2-2: 2009

#### **Summary of Sterilization and Shelf Life Discussion**

The Olympus PK Spatula is delivered in a sterile state and is intended for single patient use only. The sterilization method used is ethylene oxide and has a shelf life of three(3) years.

## **Summary of Performance Testing**

The following performance tests were conducted:

- Dimensional Measurements
- Cutting and Coagulation equivalency to predicate
- Expected forces on devices
- Design feature testing (rotation and button activation)
- Shelf Life
- Sterilization
- Biocompatibility

#### **Substantial Equivalence**

The proposed PK Spatula has the same intended use, design, and scientific technology as the Predicate PlasmaCision Laparoscopic Spatula (K041633). Both devices are of the same design, intended for the same patient population, have the identical indications for use, and use the same scientific technology. In addition there were no new issues of safety or effectiveness found with the proposed device.

#### **Conclusion:**

In summary, the PK Spatula is substantially equivalent to the predicate device and presents no new questions of safety or efficacy.